

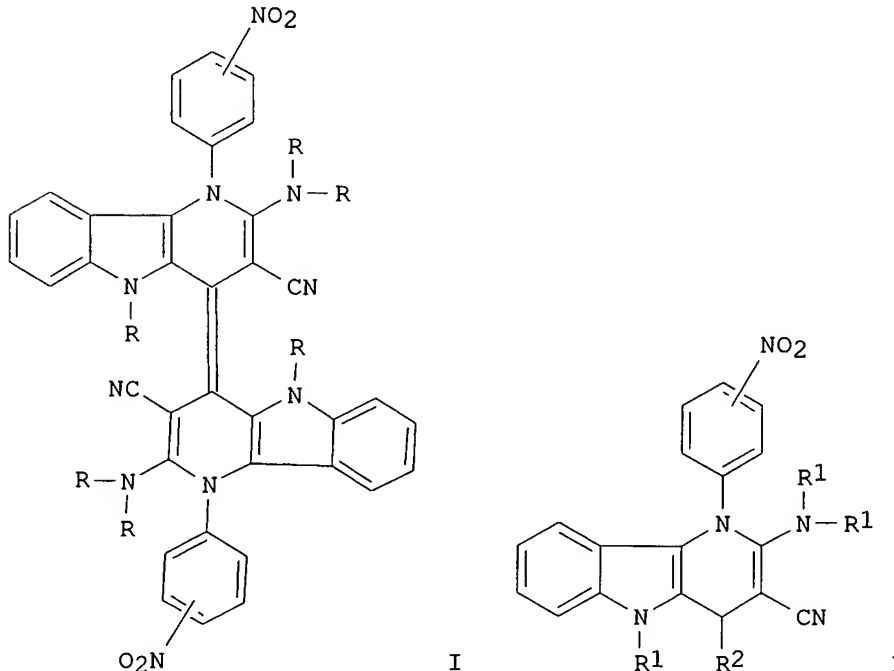
L4 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
 ACCESSION NUMBER: 2002:539680 CAPLUS
 DOCUMENT NUMBER: 137:93737
 TITLE: Preparation of pyridoindoles as anti-AIDS agents
 INVENTOR(S): Rice, William G.; Huang, Mingjun; Buckheit, Robert W., Jr.; Covell, David G.; Czerwinski, Grzegorz; Michejda, Christopher J.
 PATENT ASSIGNEE(S): The Government of the United States of America, Secretary of Health and Human Services, USA; Makarov, Vadim
 SOURCE: PCT Int. Appl., 49 pp.
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002055520	A2	20020718	WO 2001-US48310	20011213
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
US 2002182151	A1	20021205	US 2001-17323	20011213

PRIORITY APPLN. INFO.: US 2000-256556P P 20001218

OTHER SOURCE(S): MARPAT 137:93737

GI



AB The title benzoylalkylindolepyridinium (BAIP) [sic] compds. I and II [wherein R and R1 = independently H or aliph.; R2 = CH₂COCH₃] were prep'd. and tested for antiviral activity against several retroviruses. I inhibit the reverse transcriptase enzymes of several retroviruses, including human immunodeficiency virus (HIV). For example, deacylation of 3-(p-nitrophenylamino)indole (80%), followed by formylaton (96%) and condensation with malonitrile (80%), afforded the (aminoindolylmethylidenyl)malononitrile intermediate. Cyclization to the 2-imino-1,2-dihydro-5H-pyrido[3,2-b]indole (60%). Methylation with MeI in acetone in the presence of anhyd. K₂CO₃ produced the unexpected 2-oxopropyl product I (R1 = Me; R2 = CH₂COCH₃; p-nitrophenyl) (III). The latter exerted antiretroviral activity against HIV-1RF, HIV-2ROD, and SIV in a std. screening cytoprotection assay with EC₅₀ values of 0.1 .mu.M, 4.79 .mu.M, and 5.65 .mu.M, resp., and CC₅₀ values > 200 .mu.M. Further studies demonstrated that III acts during the late phase of infection, after the provirus has integrated into the host cell genome, and that cells treated with III showed reduced virion-assocd. reverse transcriptase activity and viral infectivity levels. I and II are useful for therapy to individuals already carrying HIV-1 variants that are resistant to AZT or classical non-nucleoside reverse transcriptase inhibitors (no data).

=> E RICE WILLIAM G/AU 25
E1 3 RICE WILLIAM D/AU
E2 4 RICE WILLIAM E/AU
E3 57 --> RICE WILLIAM G/AU
E4 1 RICE WILLIAM GLENN/AU
E5 2 RICE WILLIAM H/AU
E6 13 RICE WILLIAM J/AU
E7 1 RICE WILLIAM JAMES/AU
E8 1 RICE WILLIAM JOHN/AU
E9 8 RICE WILLIAM L/AU
E10 7 RICE WILLIAM L R/AU
E11 4 RICE WILLIAM M/AU
E12 9 RICE WILLIAM R/AU
E13 2 RICE WILLIAM T/AU
E14 7 RICE WILLIAM Y JR/AU
E15 1 RICE WILLIAM YATES JR/AU
E16 1 RICE WIN E/AU
E17 1 RICE WINSTON/AU
E18 2 RICE WM/AU
E19 2 RICE WM E/AU
E20 1 RICE WM H/AU
E21 2 RICE WM J/AU
E22 2 RICE WM L R/AU
E23 1 RICE WM N/AU
E24 1 RICE WM S/AU
E25 1 RICE WM T/AU

=> S (E3 OR E4) AND (ANTIVIRAL)
57 "RICE WILLIAM G"/AU
1 "RICE WILLIAM GLENN"/AU
37234 ANTIVIRAL
766 ANTIVIRALS
37389 ANTIVIRAL
(ANTIVIRAL OR ANTIVIRALS)
L1 41 ("RICE WILLIAM G"/AU OR "RICE WILLIAM GLENN"/AU) AND (ANTIVIRAL)

=> S (E3 OR E4) AND (HIV)
57 "RICE WILLIAM G"/AU
1 "RICE WILLIAM GLENN"/AU
47636 HIV
80 HIVS
47643 HIV
(HIV OR HIVS)
L2 44 ("RICE WILLIAM G"/AU OR "RICE WILLIAM GLENN"/AU) AND (HIV)

=> S (E3 OR E4) AND (BAIP)
57 "RICE WILLIAM G"/AU
1 "RICE WILLIAM GLENN"/AU
3 BAIP
L3 1 ("RICE WILLIAM G"/AU OR "RICE WILLIAM GLENN"/AU) AND (BAIP)

=> d 13 ibib abs

L3 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 2002:539680 CAPLUS
DOCUMENT NUMBER: 137:93737
TITLE: Preparation of pyridoindoles as anti-AIDS agents
INVENTOR(S): Rice, William G.; Huang, Mingjun; Buckheit,
Robert W., Jr.; Covell, David G.; Czerwinski,
Grzegorz; Michejda, Christopher J.
PATENT ASSIGNEE(S): The Government of the United States of America,
Secretary of Health and Human Services, USA; Makarov,

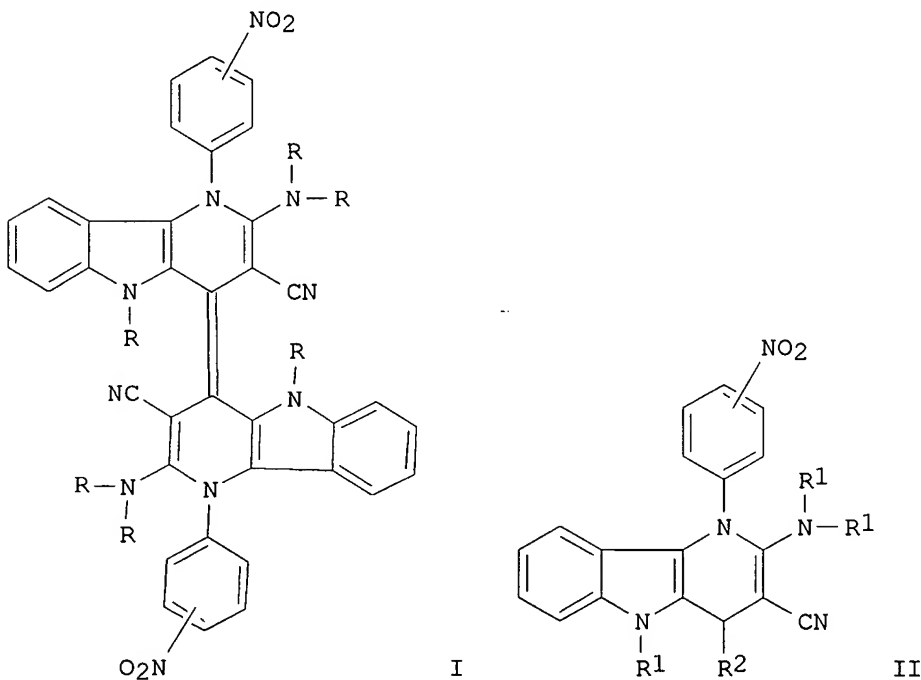
SOURCE: Vadim
PCT Int. Appl., 49 pp.
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002055520	A2	20020718	WO 2001-US48310	20011213
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GO, GW, ML, MR, NE, SN, TD, TG				
US 2002182151	A1	20021205	US 2001-17323	20011213

PRIORITY APPLN. INFO.: US 2000-256556P P 20001218

OTHER SOURCE(S): MARPAT 137:93737

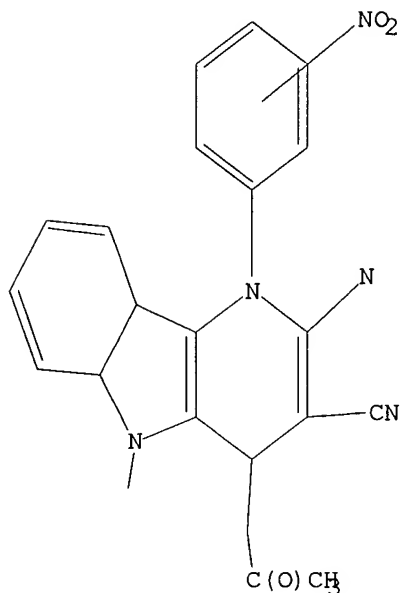
GI



AB The title benzoylalkylindolepyridinium (**BAIP**) [sic] compds. I and II [wherein R and R1 = independently H or aliph.; R2 = CH₂COCH₃] were prep'd. and tested for antiviral activity against several retroviruses. I inhibit the reverse transcriptase enzymes of several retroviruses, including human immunodeficiency virus (HIV). For example, deacylation of 3-(p-nitrophenylamino)indole (80%), followed by formylation (96%) and condensation with malonitrile (80%), afforded the (aminoindolylmethylidenyl)malononitrile intermediate. Cyclization to the 2-imino-1,2-dihydro-5H-pyrido[3,2-b]indole (60%). Methylation with MeI in

acetone in the presence of anhyd. K₂CO₃ produced the unexpected 2-oxopropyl product I (R₁ = Me; R₂ = CH₂COCH₃; p-nitrophenyl) (III). The latter exerted antiretroviral activity against HIV-1RF, HIV-2ROD, and SIV in a std. screening cytoprotection assay with EC₅₀ values of 0.1 .μ.M, 4.79 .μ.M, and 5.65 .μ.M, resp., and CC₅₀ values > 200 .μ.M. Further studies demonstrated that III acts during the late phase of infection, after the provirus has integrated into the host cell genome, and that cells treated with III showed reduced virion-assocd. reverse transcriptase activity and viral infectivity levels. I and II are useful for therapy to individuals already carrying HIV-1 variants that are resistant to AZT or classical non-nucleoside reverse transcriptase inhibitors (no data).

4 HAS NO ANSWERS
L4 STR



Structure attributes must be viewed using STN Express query preparation.

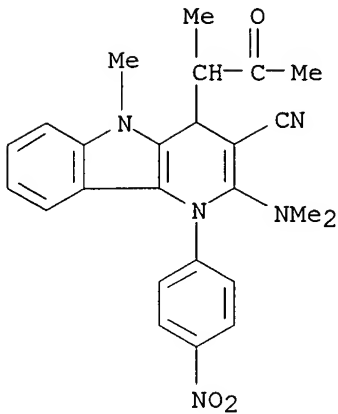
=> s 14 full
FULL SEARCH INITIATED 12:46:05 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 19 TO ITERATE

100.0% PROCESSED 19 ITERATIONS 2 ANSWERS
SEARCH TIME: 00.00.01

L5 2 SEA SSS FUL L4

=> d 15 1-2

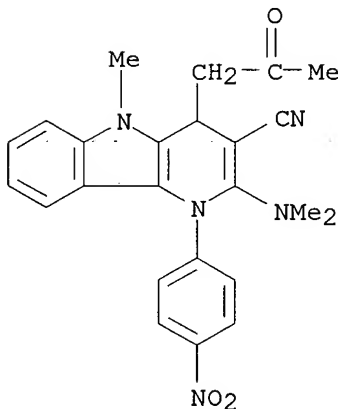
L5 ANSWER 1 OF 2 REGISTRY COPYRIGHT 2003 ACS
RN 184772-01-8 REGISTRY
CN 1H-Pyrido[3,2-b]indole-3-carbonitrile, 2-(dimethylamino)-4,5-dihydro-5-methyl-4-(1-methyl-2-oxopropyl)-1-(4-nitrophenyl)- (9CI) (CA INDEX NAME)
FS 3D CONCORD
MF C25 H25 N5 O3
SR CA
LC STN Files: CA, CAPLUS



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1962 TO DATE)
 1 REFERENCES IN FILE CAPLUS (1962 TO DATE)

L5 ANSWER 2 OF 2 REGISTRY COPYRIGHT 2003 ACS
 RN 167954-18-9 REGISTRY
 CN 1H-Pyrido[3,2-b]indole-3-carbonitrile, 2-(dimethylamino)-4,5-dihydro-5-methyl-1-(4-nitrophenyl)-4-(2-oxopropyl)- (9CI) (CA INDEX NAME)
 FS 3D CONCORD
 MF C24 H23 N5 O3
 SR CA
 LC STN Files: CA, CAPLUS, CHEMCATS, TOXCENTER, USPATFULL



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

4 REFERENCES IN FILE CA (1962 TO DATE)
 4 REFERENCES IN FILE CAPLUS (1962 TO DATE)

=> file caplus		
COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	151.51	180.89
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE ENTRY	TOTAL SESSION

CA SUBSCRIBER PRICE

0.00

-0.65

FILE 'CAPLUS' ENTERED AT 12:46:39 ON 05 FEB 2003
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
COPYRIGHT (C) 2003 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications. The CA Lexicon is the copyrighted intellectual property of the American Chemical Society and is provided to assist you in searching databases on STN. Any dissemination, distribution, copying, or storing of this information, without the prior written consent of CAS, is strictly prohibited.

FILE COVERS 1907 - 5 Feb 2003 VOL 138 ISS 6
FILE LAST UPDATED: 4 Feb 2003 (20030204/ED)

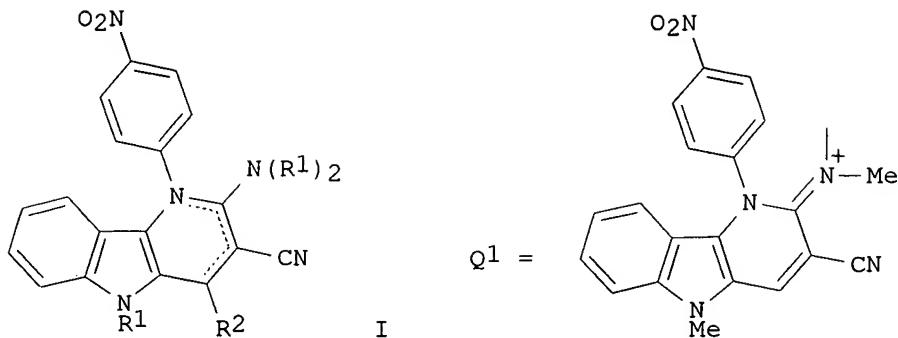
This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s 15
L6 4 L5

=> d 16 1-4 ibib abs hitstr

L6 ANSWER 1 OF 4 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 2002:575081 CAPLUS
DOCUMENT NUMBER: 137:125149
TITLE: Preparation of pyridoindoles as reverse transcriptase inhibitors.
INVENTOR(S): Rice, William G.; Huang, Mingjun; Buckheit, Robert W., Jr.; Covell, David G.; Czerwinski, Grzegorz; Michejda, Christopher J.
PATENT ASSIGNEE(S): The Government of the United States of America, Department of Health and Human Services, USA
SOURCE: PCT Int. Appl., 62 pp.
CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002059123	A2	20020801	WO 2001-US48311	20011213
W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM			
RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			
PRIORITY APPLN. INFO.:			US 2000-256581P	P 20001218
OTHER SOURCE(S):			MARPAT 137:125149	



AB Title compds. (I; R1 = alkyl; R2 = H, alkylamide, Q1; dotted lines = optional double bonds), were prepd. Thus, 1-(4-nitrophenyl)-2-methylimino-3-cyano-5-methyl-1,2-dihydro-5H-pyrido[3,2-b]indole (prepn. given) was refluxed with K₂CO₃, MeI, and acetone for 45 h to give 1-(4-nitrophenyl)-2-dimethylamino-3-cyano-4-(2-oxopropyl)-5-methyl-1,2-dihydro-5H-pyrido[3,2-b]indole. The latter showed IC₅₀ = 0.1 μ M against HIV-1 RF in CEM-SS cells.

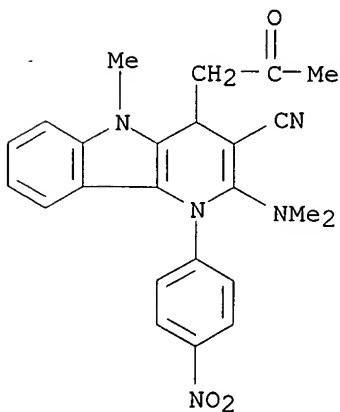
IT 167954-18-9P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of pyridoindoles as reverse transcriptase inhibitors)

RN 167954-18-9 CAPLUS

CN 1H-Pyrido[3,2-b]indole-3-carbonitrile, 2-(dimethylamino)-4,5-dihydro-5-methyl-1-(4-nitrophenyl)-4-(2-oxopropyl)- (9CI) (CA INDEX NAME)



L6 ANSWER 2 OF 4 CAPLUS COPYRIGHT 2003 ACS

ACCESSION NUMBER: 2002:539680 CAPLUS

DOCUMENT NUMBER: 137:93737

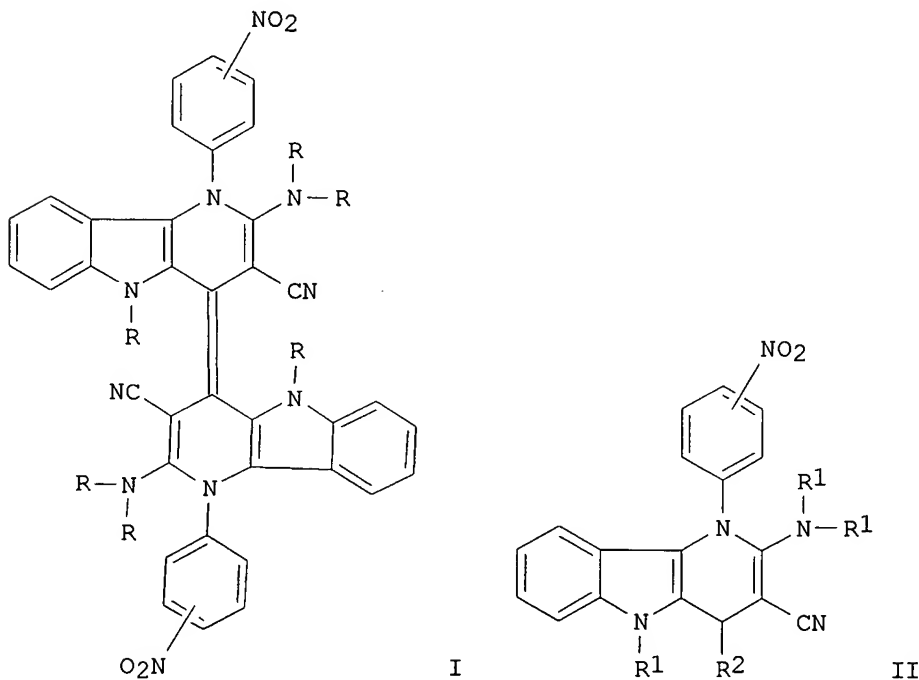
TITLE: Preparation of pyridoindoles as anti-AIDS agents

INVENTOR(S): Rice, William G.; Huang, Mingjun; Buckheit, Robert W., Jr.; Covell, David G.; Czerwinski, Grzegorz; Michejda, Christopher J.

PATENT ASSIGNEE(S): The Government of the United States of America, Secretary of Health and Human Services, USA; Makarov, Vadim

SOURCE: PCT Int. Appl., 49 pp.
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION: CODEN: PIXXD2

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002055520	A2	20020718	WO 2001-US48310	20011213
W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM			
RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			
US 2002182151	A1	20021205	US 2001-17323	20011213
PRIORITY APPLN. INFO.:			US 2000-256556P	P 20001218
OTHER SOURCE(S):			MARPAT 137:93737	
CJ				



AB The title benzoylalkylindolepyridinium (BAIP) [sic] compds. I and II [wherein R and R1 = independently H or aliph.; R2 = CH2COCH3] were prep'd. and tested for antiviral activity against several retroviruses. I inhibit the reverse transcriptase enzymes of several retroviruses, including human immunodeficiency virus (HIV). For example, deacylation of 3-(p-nitrophenylamino)indole (80%), followed by formylation (96%) and condensation with malonitrile (80%), afforded the (aminoindolylmethylidenyl)malononitrile intermediate. Cyclization to the

2-imino-1,2-dihydro-5H-pyrido[3,2-b]indole (60%). Methylation with MeI in acetone in the presence of anhyd. K2CO3 produced the unexpected 2-oxopropyl product I (R1 = Me; R2 = CH2COCH3; p-nitrophenyl) (III). The latter exerted antiretroviral activity against HIV-1RF, HIV-2ROD, and SIV in a std. screening cytoprotection assay with EC50 values of 0.1 .mu.M, 4.79 .mu.M, and 5.65 .mu.M, resp., and CC50 values > 200 .mu.M. Further studies demonstrated that III acts during the late phase of infection, after the provirus has integrated into the host cell genome, and that cells treated with III showed reduced virion-assocd. reverse transcriptase activity and viral infectivity levels. I and II are useful for therapy to individuals already carrying HIV-1 variants that are resistant to AZT or classical non-nucleoside reverse transcriptase inhibitors (no data).

IT

167954-18-9P

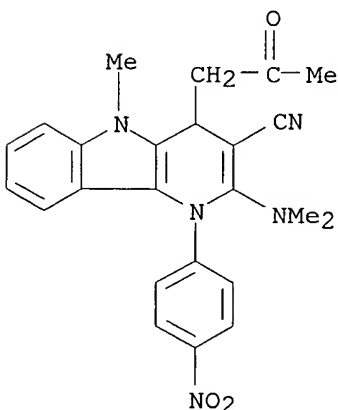
RL: ADV (Adverse effect, including toxicity); PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 (antiretroviral agent; prepn. of pyridoindole anti-AIDS agents via cyclization and subsequent derivatization of (aminoindolylmethylidenyl)malononitrile)

RN

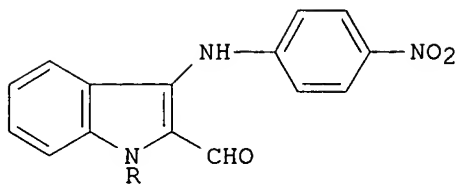
167954-18-9 CAPLUS

CN

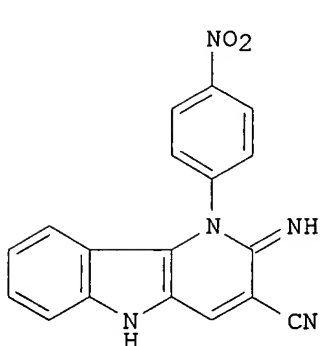
1H-Pyrido[3,2-b]indole-3-carbonitrile, 2-(dimethylamino)-4,5-dihydro-5-methyl-1-(4-nitrophenyl)-4-(2-oxopropyl)- (9CI) (CA INDEX NAME)



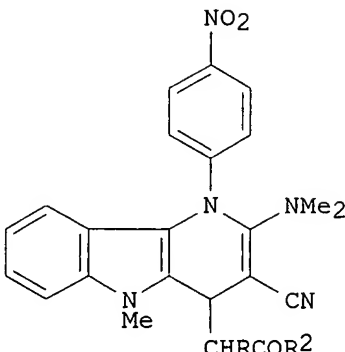
L6 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2003 ACS
 ACCESSION NUMBER: 1996:735136 CAPLUS
 DOCUMENT NUMBER: 126:47132
 TITLE: 2-Formyl-3-(arylarnino)indoles in the synthesis of 1,2- and 1,4-dihydro-5H-pyrido[3,2-b]indoles (dihydro-.delta.-carbolines)
 AUTHOR(S): Ryabova, S. Yu.; Alekseeva, L. M.; Granik, V. G.
 CORPORATE SOURCE: TSKHLS, VNIKHFI, Moscow, Russia
 SOURCE: Khimiko-Farmatsevticheskii Zhurnal (1996), 30(9), 29-34
 PUBLISHER: Izdatel'stvo Folium
 DOCUMENT TYPE: Journal
 LANGUAGE: Russian
 GI



I



II



III

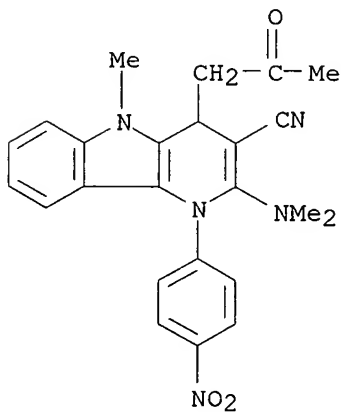
AB Condensation of the CHO group of title aldehydes I ($R = H, Ac$) with malononitrile, followed by cyclization, gave 1,2-dihydro- δ -carboline II, which reacted with MeI and carbonyl compds. to give 1,4-dihydro- δ -carbolines III [$R = H, R1 = Me; RR1 = (CH2)4; R = R1 = Me$].

IT **167954-18-9P 184772-01-8P**

RL: SPN (Synthetic preparation); PREP (Preparation)
(prepn. of)

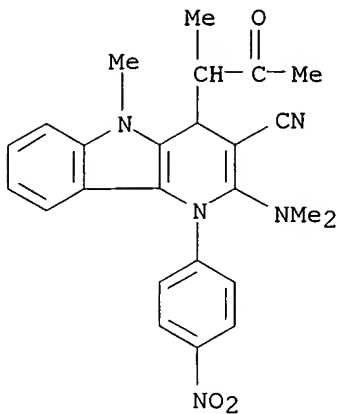
RN 167954-18-9 CAPLUS

CN 1H-Pyrido[3,2-b]indole-3-carbonitrile, 2-(dimethylamino)-4,5-dihydro-5-methyl-1-(4-nitrophenyl)-4-(2-oxopropyl)- (9CI) (CA INDEX NAME)

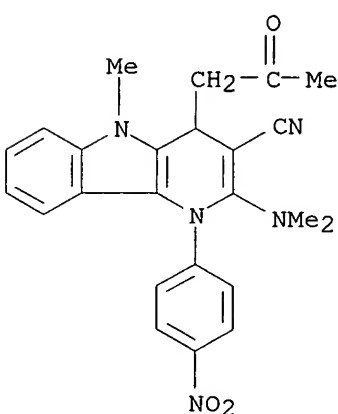


RN 184772-01-8 CAPLUS

CN 1H-Pyrido[3,2-b]indole-3-carbonitrile, 2-(dimethylamino)-4,5-dihydro-5-methyl-4-(1-methyl-2-oxopropyl)-1-(4-nitrophenyl)- (9CI) (CA INDEX NAME)



L6 ANSWER 4 OF 4 CAPLUS COPYRIGHT 2003 ACS
 ACCESSION NUMBER: 1995:622244 CAPLUS
 DOCUMENT NUMBER: 123:198747
 TITLE: A new approach to the synthesis of 1,2- and 1,4-dihydropyrido[3,2-b]indole derivatives
 AUTHOR(S): Ryabova, Svetlana Yu.; Alekseeva, Lyudmila M.; Granik, Vladimir G.
 CORPORATE SOURCE: Cent. Medicinal Chem., All-Russian Res. Chem.-Pharmaceutical Inst., Moscow, 119815, Russia
 SOURCE: Mendeleev Communications (1995), (3), 107-9
 CODEN: MENCEX; ISSN: 0959-9436
 PUBLISHER: Russian Academy of Sciences
 DOCUMENT TYPE: Journal
 LANGUAGE: English
 OTHER SOURCE(S): CASREACT 123:198747
 AB Intramol. cyclization of .alpha.-cyano-.beta.- (3-p-nitrophenylaminoindol-2-yl)acrylonitrile yields 1-p-nitrophenyl-2-imino-3-cyano-1,2-dihydropyrido[3,2-b]indole, methylation of which by Me iodide in acetone in the presence of potassium carbonate is accompanied by the addn. of acetonyl anion and formation of 1-p-nitrophenyl-2-dimethylamino-3-cyano-4-acetonyl-1,4-dihydropyrido[3,2-b]indole.
 IT 167954-18-9P
 RL: SPN (Synthetic preparation); PREP (Preparation)
 (prep. of dihydropyrido[3,2-b]indoles)
 RN 167954-18-9 CAPLUS
 CN 1H-Pyrido[3,2-b]indole-3-carbonitrile, 2-(dimethylamino)-4,5-dihydro-5-methyl-1-(4-nitrophenyl)-4-(2-oxopropyl)- (9CI) (CA INDEX NAME)



→ LIBRARY